





International Journal of Hydrogen Energy 000 (2000) 000-000

www.elsevier.com/locate/ijhydene

## Synthesis and characterization of novel hydride compounds

Randell L. Mills\*, Bala Dhandapani, Mark Nansteel, Jiliang He, Tina Shannon, Alex Echezuria

BlackLight Power, Inc., 493 Old Trenton Road, Cranbury, NJ 08512, USA

#### Abstract

Novel inorganic alkali and alkaline earth hydrides of the formula MHX and MHMX wherein M is the metal, X, is a singly negatively charged anion, and H comprises a novel high binding energy hydride ion were synthesized in which temperature gas cell by reaction of atomic hydrogen with a catalyst and MX or MX2 corresponding to an alkali metal or alkaline earth metal compound, respectively. Novel hydride compounds were identified by time of flight secondary ion mass spectroscopy, X-ray photoelectron spectroscopy, proton nuclear magnetic resonance spectroscopy, and thermal decomposition with analysis by gas chromatography, and mass spectroscopy. © 2000 International Association for Hydrogen Energy. Published by Elsevier Science Ltd. All rights reserved.

## 1. Introduction

Typically the emission of extreme ultraviolet light from hydrogen gas is achieved via a discharge at high voltage, a high power inductively coupled plasma, or a plasma created and heated to extreme temperatures by RF coupling (e.g.  $> 10^6$  K) with confinement provided by a toroidal magnetic field. Intense extreme ultraviolet (EUV) emission was observed at low temperatures (e.g.  $\approx 10^3$  K) from hydrogen and certain atomized elements on that gaseous ions which ionize at integer multiples of the example, strontium ionizes at integer multiples of the example, strontium ionizes at integer multiples of the potential energy of atomic hydrogen. Intense EUV hydrogen plasma emission was observed at low temperatures ( $\approx \approx 10^3$  K) when atomic hydrogen was generated at a lungsten filament that heated a titanium dissociator and atomic strontium was vaporized from the metal by heating No emission was observed when sodium, magnesium, or barium replaced strontium or when argon replaced hydrogen with strontium. Furthermore, the power balance of a gas cell having atomized hydrogen and strontium was ineasured by integrating the total light

• Corresponding author. Tel.: +1-609-490-1040; fax: +1-609-490-1066.

E-mail address: rmills@blacklightpower.com (R.L. Mills).

output corrected for detector response and energy over the visible range [6]. A control cell was identical except that sodium eplaced strontium. In this case, over 4000 times the power of the strontium cell was required in order to aelfieve that same optically measured light output power. A plasma formed at a cell voltage of about 250 V in the cell with hydrogen alone and in the cell with hydrogen and sodium; whereas, a plasma formed in the strontium cell at the extremely low voltage of about 2 V.

Based on their exceptional emission, we used potassium, cesium, rubidium, calcium, and strontium metals as catalysts to release energy from atomic hydrogen to form novel compounds [1,2]. The theory is given in the Appendix.

Novel inorganic alkali and alkaline earth hydrides of the formula MHX and MHMX wherein M is the metal, X, is a singly negatively charged anion, and H comprises a novel high binding energy hydride ion were synthesized in a high-temperature gas cell by reaction of atomic hydrogen with a catalyst and MX or MX<sub>2</sub> corresponding to an alkali metal or alkaline earth metal compound, respectively. For example, atomic hydrogen was reacted with strontium vapor and SrBr<sub>2</sub> to form SrHBr. Novel hydride compounds such as SrHBr were identified by time of flight secondary ion mass spectroscopy, X-ray photoelectron spectroscopy, proton nuclear magnetic resonance spectroscopy, and thermal decomposition with analysis by gas chromatography,

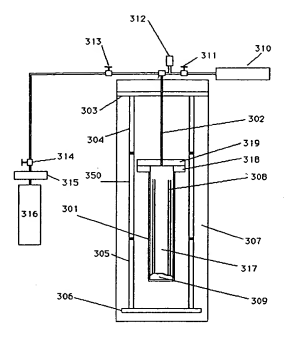


Fig. 1. Stainless steel gas cell comprising a screen dissociator, metal catalyst, and alkali or alkaline earth halide as the reactant. The components were: 301 — stainless steel cell; 317 — internal cavity of cell; 318 — high vacuum conflat flange; 319 — mating blank conflat flange; 302 — stainless steel tube vacuum line and gas supply line; 303 — lid to the kiln or top insulation, 304, 305, 350, and 306 — heaters; 307 — high temperature insulation; 308 — screen dissociator; 309 — powdered alkali or alkaline earth halide reactant; 310 — high vacuum turbo pump; 312 — pressure gauge; 311 — vacuum pump valve; 313 — valve; 314 — valve; 315 — regulator; 316 — hydrogen tank.

and mass spectroscopy. Hydride ions with increased bine ing energies form novel compounds with potential broad applications such as a high-voltage battery for consumer electronics and electric vehicles. In addition, these nove compositions of matter and associated technology machave far-reaching applications in many indistricts including chemical, electronics, computer, military, energy, and aerospace in the form of products such as mopellants, solid fuels, surface coatings, structural malerials, and chemical processes.

## 2. Experimental

## 2.1. Synthesis

# 2.1.1. Potassium indachydride, KHI, synthesis in a 3.0 l stainless steel control

Potassium iodo hydride was prepared in a stainless steel gas cell shown in Fig. 1 comprising a Ni screen hydrogen dissociator (Belleville Wire Cloth Co., Inc.), potassium metal catalyst (Aldrich Chemical Company), and KI (Aldrich Chemical Company 99.9%). The 316-stainless

steel cell was in the form of a tube having an internal cavity of 375 mm in length and 140 mm in diameter. The wall thickness was 6.35 mm. The bottom of the cell was closed by a 6.35 mm thick circular plate of 316 stainless steel that was welded to the cylinder. The top end of the cell was welded to a bored-through 304 stainless steel conflat-type flange with 8 in nominal diameter. A mating blank flange was bolted to the bored-through flange with 20 silver-plated bolts. A flange gasket was silver-plated copper. A 1.27 cm OD tube was welded into a hole at the center of the blank flange. This tube was closed at one end and extended 20 cm into the reactor, serving as a thermowell. A 9.5 mm OD stainless tube was welded to the flange approximately 4 cm from the flange center. This tube served as the vacuum line from the cell as well as a hydrogen or helium supply line to the cell.

The reactor was heated in a 10 kW refractory brick kiln (L & L Kiln Model JD230). The kiln had thee heating zones and a heated floor that were each heated by separate radiant elements. The zone temperatures were independently controlled by a Dynatrol controlle. The reactor was instrumented with 5 type-K thermocourles. Two thermocourles were located in the central thermocourles. Three thermocourles were fixed to the external surface of the reactor and were located near the base, a mid-height, and near flange-level. The reactor was connected through bellows-type valves to a turbo vacuum pump. The vacuum level was measured by a 0-100 Torr paratron vacuum gauge. Pressures above 100 Torr were measured by standard dial-type pressure gauges. Temperature and pressure data was logged to a data acquisition system at 5 min intervals.

Approximately 290 g of nickel screen (0.5 mm wire, 2 mm lesh was placed circumferentially around the reactor inner all of the cell. 125 g of dry KI were placed in a stainless steel crucible on the reactor base. The reactor was flooded with argon gas. 1.7 g of metallic potassium was placed in a smaller stainless steel crucible and this crucible was placed in the larger one with the KI crystals. The reactor was sealed and placed in the kiln. The system was evacuated for 2.5 h. The reactor was pressurized with hydrogen gas to a pressure of 10 Torr and sealed. The kiln was heated to 650°C at the rate of 300°C/h. The reactor was held at 650°C for 72 h. Hydrogen was added to the system periodically to maintain a pressure level of 10 Torr. The reactor was then evacuated for 1 h while at 650°C. The kiln and reactor were cooled to room temperature by forced convection in about 2 h while pumping continued. At room temperature the system was filled with helium gas to a pressure of 1.3 bar. The sealed reactor was then opened. About 125 g of green crystals were observed to have formed in the stainless steel crucible.

## 2.1.2. Strontium fluoro hydride synthesis in a 40 cm³ stainless steel gas cell reactor

Strontium fluoro hydride was prepared in a stainless steel gas cell shown in Fig. 1 comprising a Ti screen hydrogen







R.L. Mills et al. | International Journal of Hydrogen Energy 000 (2000) 000-000

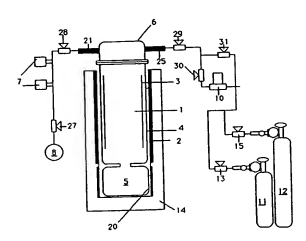


Fig. 2. Quartz gas cell comprising a Ni screen dissociator, potassium metal catalyst, and  $K_2CO_3$  as the reactant. The components were: 1— internal cavity of cell; 3— nickel hydrogen dissociator; 4— quartz tube cell with Conflat style flange; 5— catalyst reservoir, 6— Pyrex cap with an identical Conflat style flange to the tube; 25— gas inlet line; 21— gas outlet line; 11— compressed gas cylinder of ultra high purity hydrogen; 13— hydrogen control valve; 12— compressed gas cylinder of ultrahigh purity helium; 15— helium control valve; 10— mass flow controller; 30— mass flow controller valve; 29— inlet valve; 31— mass flow controller bypass valve; 8— molecular drag pump; 27— vacuum pump valve; 28— outlet valve; 7— 0 to 1000 torr Baratron pressure gauge and a 0 to 100 torr Baratron pressure gauge; 20— catalyst reservoir band heater; 14— Zircar AL-30 insulation package; 2— Mellen cell heater.

dissociator (Belleville Wire Cloth Co., Inc.), strontium metal catalyst (Alfa Aesar), and  $SrF_2$  (Aldrich Chemical Company 99.99%) as the reactant. The 304-stainless steel cell was more than 10 the form of a tube having an internal cavity of 75 mm in length and 28 mm in diameter. The top end of the cell was welded to a high vacuum  $2\frac{3}{4}$  in bored through conflat flange. The mating blank conflat flange contained a single opting in which was welded a  $\frac{1}{4}$  in diameter stainles used tube that was 10 cm in length and contained a needle value to isolate the reactor from the rest of the gas manufold spilorer plated copper gasket was placed between the two flanges. The two flanges are held together with 6 circumferential bolts. The bottom of the  $\frac{1}{4}$  in tube was flush with the bottom surface of the top flange. The  $\frac{1}{4}$  in tube gerved as a vacuum line and also as a hydrogen or helium supply line to the cell. The cell was surrounded by a claim shell heater (Mellen Company), which was in turn surrounded by insulation. The heater was controlled by warrage.

About 10-g of Ti screen, 12 g of crystalline SrF<sub>2</sub> and 0.6 g of strontium metal was added to the cell under an argon atmosphere. The cell was then continuously evacuated with the isolation needle valve open using a high vacuum turbo pump to reach 20 mTorr measured by a pressure gauge (MKS). The cell was heated by supply-

ing power to the heaters. The temperature of the cell was measured with a type K thermocouple (Omega). The cell temperature was then slowly increased to 100°C using the heaters. The vacuum pump valve was closed. Hydrogen was slowly added to maintain a pressure of 1 atm, and the needle valve was closed to isolate the reactor system. The temperature of the cell was then slowly increased to 650°C. Hydrogen was added periodically to maintain 1 atm using the needle valve. After 72 h, the temperature of the cell was reduced to room temperature. The reactor was flushed with helium and closed using the needle valve. It was then opened in an argon environment chamber to recover the strontium fluoro hydride. A white solid was obtained.

# 2.1.3. Potassium hydride potassium hydrogen carbonate synthesis in a quartz gas cell reactor

Potassium hydride potassium hydroten carbonate was prepared in a quartz gas cell shown in Fig. 2 comprising a nickel screen hydrogen dissociator (Belleville Wire Cloth Co., Inc.), potassium metal catalyst (Aldrich Chemical Company), and K<sub>2</sub>CO<sub>3</sub> (Aldrich Chemical Company), and K<sub>2</sub>

The gas was supplied to the cell through the inlet from a compressed gas cylinder of ultra high-purity hydrogen controlled by a hydrogen control valve. Helium gas was supplied to the cell through the same inlet from a compressed gas cylinder of ultrahigh purity helium controlled by helium control valve. The flow of helium and hydrogen to the cell is further controlled by a mass flow controller, a mass flow controller valve, an inlet valve, and a mass flow controller bypass valve. The bypass valve was closed during filling of the cell. Excess gas was removed through the gas outlet by a molecular drag pump capable of reaching pressures of 10-4 Torr controlled by vacuum pump valve and outlet valve. Pressures were measured by a 0-10 Torr Baratron pressure gauge. The reactor and the catalyst reservoir, were heated independently using clam shell heaters (Mellen Company) powered by Variacs. The temperature was recorded using a K-type thermocouple placed close to the quartz

The cell was operated under flow conditions with a total pressure of less than two (2) Torr of hydrogen or control helium via mass flow controller. About 20 g of K<sub>2</sub>CO<sub>3</sub> (Aldrich Chemical Company, 99.9%) was placed in the

catalyst reservoir and about 50 g, 12×12 cm of nickel screen dissociator (Belleville Wire Cloth Co., Inc.) was treated with 0.6 M K<sub>2</sub>CO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> and dried at 130°C overnight and then placed in the center of the reactor. The reactor was evacuated to about 50 mTorr and hydrogen was introduced using the mass flow controller to maintain a pressure of about 2 Torr. The temperature of the reactor was increased to about 800-850°C to facilitate atomization of hydrogen, and the catalyst reservoir temperature was increased to 850°C to vaporize the catalyst. The reaction started when the catalyst vapor reached the hot zone of the reactor. The reaction was allowed to continue for about 120 h. The reaction was terminated by cooling the reactor and the catalyst reservoir to room temperature in hydrogen. The system was purged and back filled with helium and sealed. The reactor was opened in an argon environmental chamber and the samples were collected and analyzed. The compound was a white powder.

## 2.1.4. Synthesis of alkali halido hydrides and alkaline earth halido hydrides

A series of alkali and alkaline earth halido hydrides (KHF, KHCl, KHBr, KHI, RbHF, RbHCl, RbHBr, RbHI, CsHF, CsHCl, CsHBr, CsHI, CaHCl, CaHBr, CaHl, SrHF, SrHCl, and SrHBr) were synthesized in gas cells as described in the previous A-B Sections with the exception that the alkali or alkaline earth metal catalyst (rubidium metal (which is a catalyst as a hydride having Rb+) and potassium, cesium, calcium, and strontium metals) corresponded to the alkali or alkaline earth halide of the product alkali or alkaline earth halido hydride. RbHF was synthesized by the catalysis of atomic hydrogen with potassium metal catalyst followed by reaction with RbF wherein the hydrogen dissociator was a nickel screen. Reactants to form hydrides of these inorganic compounds obtained from Alfa Asak were KF (99.9%), KCl (ACS grade 99+(99.9%) Rbl (99.9%), CsF (99.9%), CsCl (99.9%), CsBr (99.2%), CsL (99.9%), CaCl<sub>2</sub> (99.9%), CaBr<sub>2</sub> (99.9%), Cal<sub>2</sub> (99.9%) SrF<sub>2</sub> (99.9%), SrCl<sub>2</sub> (99.9%), and SrBr<sub>2</sub> (99.9%). If the analytical analyses, each starting compound as a control.

## 2.2. ToF-SIMS characterization

The crystalline samples were spirnkled onto the surface of a double-sided adhesive tape and characterized using a Physical Electronics TFS 1000 ToF-SIMS instrument. The primary ion gun utilized a 69 Ga+ liquid metal source. In order to remove surface contaminants and expose a fresh surface, the samples were sputter cleaned for 30 s using a 40 µm x 10 µm raster. The aperture setting was 3, and the ion cultern was 600 pA resulting in a total ion dose of 101 ions/cm<sup>2</sup>

During acquisition, the ion gun was operated using a bunched (pulse width 4 ns bunched to 1 ns) 15 kV beam

[7,8]. <sup>1</sup> The total ion dose was  $10^{12}$ ions/cm<sup>2</sup>. Charge neutralization was active, and the post accelerating voltage was 8000 V. Three different regions on each sample of  $(12 \mu m)^2$ ,  $(18 \mu m)^2$ , and  $(25 \mu m)^2$  were analyzed. The positive and negative SIMS spectra were acquired. Representative post sputtering data is reported.

#### 2.3. XPS characterization

A series of XPS analyses were made on the crystalline samples of KHCl and KCl using a Scienta 300 XPS Spectrometer. The fixed analyzer transmission mode and the sweep acquisition mode were used. The step energy in the survey scan was 0.5 eV, and the step energy in the high resolution scan was 0.15 eV. In the survey scan, the time per step was 0.4 s, and the number of sweeps was 4. In the high resolution scan, the time per step was 0.3 s, and the number of sweeps was 30. C 1s at 284.5 eV was used as the internal standard.

The binding energies and features of conference electrons of a series of alkali and alkaline earth naildo in drides (KHCl, KHI, RbHCl, RbHI, CsHBr, CaHCl, CaHBr, CaHI, SrHF, SrHCl, and SrHBr) were analyzed by APS. XPS analysis was conducted on a Kratos XSAM-800 spectrometer using nonmonochromatic Alka (1468.6 eV) radiation. Samples were crushed in a glove box under argon and mounted on an analysis stub with copper tape. A piece of gold foil was stuck into the sample for calibration. The samples were transferred under an inert atmosphere. A survey spectrum was run from 1000 to 0 eV. For quantitative analysis, high resolution spectra were run on core level electrons of interstrench as the Rb3d and Cs3d electrons. For KI and KHI, a high third spectrum of the low binding energy region was also run from 100 to 0 eV that corresponded to the sur-Spectrum. Fixed analyzer transmission (FAT) mode was used in all measurements. For the survey scan, a pass energy of 320 eV was employed. A pass energy of 40 eV was used for high resolution scans. In the cases were a charging effect was observed, the spectrum was corrected by using a calibration of the effect with the Au4f<sub>7/2</sub> peak at 84.0 eV as a first standard and the C1s peak at 284.6 eV as a second standard.

## 2.4. NMR spectroscopy

<sup>1</sup>H MAS NMR was performed on solid samples of KHI, KHCl, KHBr, and RbHF. The data was obtained on a custom built spectrometer operating with a Nicolet 1280 computer. Final pulse generation was from a tuned Henry radio amplifier. The <sup>1</sup>H NMR frequency was 270.6196 MHz. A 5  $\mu$  s pulse corresponding to a 41° pulse length and a 3 recycle delay were used. The window was ±20 kHz. The spin speed was 4.0 kHz. (The spin speed was varied to confirm real peaks versus side bands. The latter changed

<sup>1</sup> For recent specifications see Ref. [8].

position with spin speed, the former were independent of spin speed.) The number of scans was 600. The offset was 1541.6 Hz, and the magnetic flux was 6.357 T. The samples were handled under a nitrogen atmosphere. Chemical shifts were referenced to external tetramethylsilane (TMS). The reference of KHI comprised KH (Aldrich Chemical Company 99%) and equivalent molar mixtures of KH (Aldrich Chemical Company 99%) and KI (Aldrich Chemical Company 99.99%) prepared in a glove box under argon. The reference of KHCl comprised KH (Aldrich Chemical Company 99%) and equivalent molar mixtures of KH (Aldrich Chemical Company 99%) and KCI (Aldrich Chemical Company 99.99%) prepared in a glove box under argon. The reference of KHBr comprised KH (Aldrich Chemical Company 99%) and equivalent molar mixtures of KH (Aldrich Chemical Company 99%) and KBr (Aldrich Chemical Company 99.99%) prepared in a glove box under argon. The reference of RbHF comprised RbH (Aldrich Chemical Company 99%).

<sup>1</sup>H MAS NMR was performed on a solid samples of SrHBr. The data were recorded on a Bruker DSX-300 spectrometer at 300.132 MHz. Samples were packed and sealed in 5 mm diameter NMR tubes under an inert atmosphere. The <sup>1</sup>H MAS NMR of strontium bromo hydride was run static. The MAS frequency was 4.1 kHz. During data acquisition, the 90° pulse length for a single pulse <sup>1</sup>H excitation was 3.4 μs; the sweep width was 147.058 kHz; the dwell time was 5.5 μs, and the acquisition time was 0.0139764 s/scan. The number of scans was typically 32.

The hydrogen composition of the samples was quantified by integrating the signal under each peak. In each case, the signal from an empty tube was baseline subtracted from the integral for each peak. The integration was calibrated using a polyethylene standard. From the sample weight, the wt% hydrogen was determined. The chemical shifts of the spectra were calibrated based on the chemical shift of HO. The accuracy was determined to be within 0.1 ppm some samples were broadened due to  ${}^{1}H_{-}{}^{1}H$  dipole interaction. The broadening from a neighboring proton interaction and the proton of the protons of the protons to a neighboring protons  ${}^{1}H_{-}{}^{2}H_{-}{}^{3}$ 

# 2.5. Thermal decomposition with analysis by gas Chromatography

Solid samples were decomposed at high temperature and quantitatively analyzed for hydrogen using gas chromatography. On Prof. ample was placed in a thermal decomposition reactor under an argon atmosphere. The reactor comprised a 174" OD by 3" long quartz tube that was sealed at one end and connected at the open end with Swagelock things to a T. One end of the T was connected to a needle valve and a HOVAC molecular drag pump. The other end

was attached to a septum port. The apparatus was evacuated to between 25 and 50 m Torr. The needle valve was closed to form a gas tight reactor. The sample was heated in the evacuated quartz chamber containing the sample with an external Nichrome wire heater using a Variac transformer. The sample was heated to above 600°C by varying the transformer voltage supplied to the Nichrome heater until the sample melted. Gas released from the sample was collected with a 500 µl gas tight syringe through the septum port and immediately injected into the gas chromatograph.

Gas samples were analyzed with a Hewlett Packard 5890 Series II gas chromatograph equipped with a thermal conductivity detector and a 60 m, 0.32 mm ID fused silica Rt-Alumina capillary PLOT column (Restek, Bellefonte, PA). The column was conditioned at 200° C for 18–72 h before each series of runs. Samples were run at -196° C using Ne as the carrier gas. The 60 m column was run with the carrier gas at 3.4 psi with the following flow rates: carrier -2.0 ml/min, auxiliary -3.4 ml/min and reference -3.5 ml/min, for a total flow rate of 89 ml/min. The split rate was 10.0 ml/min.

The control hydrogen gas was ultrahign purity (MG Industries). Control samples were also treated by the same method as the samples of the novel compounds.

# 2.6. Thermal decomposition with analysis by mass spectroscopy

Mass spectroscopy was performed on the gases released from the thermal decomposition of the samples. One end of mic ID fritted capillary tube containing about 5 mg Sample was sealed with a 0.25 in. Swagelock union and hug (Spelock Co., Solon, OH). The other end was conpected directly to the sampling port of a Dycor System 1000 drapole Mass Spectrometer (Model D200MP, Ametek, Inc., Pittsburgh, PA with a HOVAC Dri-2 Turbo 60 Vacuum System). The capillary was heated with a Nichrome wire heater wrapped around the capillary. The mass spectrum was obtained at the ionization energy of 70 and 30 eV at different sample temperatures in the region m/e = 0-50. With the detection of hydrogen indicated by a m/e = 2 peak, the intensity as a function of time for masses m/e = 1, m/e = 2, m/e = 3, m/e = 4, m/e = 5, and m/e = 18 was obtained while changing the ionization potential (IP) of the mass spectrometer from 30 to 70 eV.

The control hydrogen gas was ultrahigh purity (MG Industries).

## 3. Results and discussion

## 3.1. ToF-SIMS

## 3.1.1. ToF-SIMS of potassium iodo hydride sample

The positive ToF-SIMS spectrum (m/e = 0-140) of KHI, the positive ToF-SIMS spectrum (m/e = 0-140) of KI, the negative ToF-SIMS spectrum (m/e = 0-140) of KHI, and

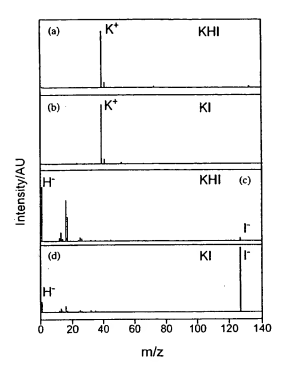


Fig. 3. (a) The positive ToF-SIMS spectrum (m/e = 0-140) of KHl. (b) The positive ToF-SIMS spectrum (m/e = 0-140) of Kl. (c) The negative ToF-SIMS spectrum (m/e = 0-140) of KHl. (d) The negative ToF-SIMS spectrum (m/e = 0-140) of Kl.

the negative ToF-SIMS spectrum (m/e = 0-140) of KI are shown in Figs. 3a-d, respectively. The positive ion spectrum of KHl and that of the KI control were dominated by the K<sup>+</sup> ion. A K<sup>2+</sup> ion was only observed in the positive ion spectrum of the KHl.

 $Ga^+$  m/z = 69,  $K_2^+$  m/z = 78,  $K(KC1)^+$  m/z (13),  $I^+$  m/z = 127,  $KI^+$  m/z = 166, and a series of positive ion  $K[K1]_n^+$  m/z = (39 + 166n) were also observed the negative ion ToF-SIMS of KHI was dominated by H with a smaller  $I^-$  peak. lodide alone dominated the negative ion ToF-SIMS of the KI control. For both,  $O^-$  m/z = 160,  $OH^-$  m/z = 17,  $CI^-$  m/z = 35,  $KI^-$  m/z = 1662 a series of negative ions  $I[K1]_n^-$  m/z = (127 + 166n) were also observed.

# 3.1.2. ToF-SIMS of polassium hydride potassium hydrogen carbonate sample

The positive 201 SIMS spectrum obtained from the KHCO<sub>3</sub> control is shown in Figs. 4 and 5. (KHCO<sub>3</sub> was used as a control versus  $205 \text{ Ge}_3$  because the former is more conservative in that it contains a larger source of H). In addition, the positive 205 For SIMS of the KHKHCO<sub>3</sub> sample is shown in Figs. 6 and 205 For SIMS hooth the control and KHKHCO<sub>3</sub> samples, the positive ion spectrum are dominated by the K + ion. Two series of positive ions  $205 \text{ K} \text{ K} \text{ K} \text{ CO}_3$ , 205 For SIMS and  $205 \text{ K} \text{ CO}_3$ , 205 For SIMS are observed

in the KHCO<sub>3</sub> control. Other peaks containing potassium include KC<sup>+</sup>,  $K_xO_y^+$ ,  $K_xO_y^+$ ,  $K_xO_y^+$ ,  $K_xO_y^+$ ,  $K_xO_y^+$ , and  $K_2^+$ . However, in the KHKHCO<sub>3</sub> sample, three new series of positive ions are observed at  $\{K[KHKHCO_3]_n^+ \ m/z = (39 + 140n), K_2OH[KHKHCO_3]_n^+ \ m/z = (95 + 140n),$  and  $K_3O[KHKHCO_3]_n^+ \ m/z = (133 + 140n)\}$ . These ions correspond to inorganic clusters containing novel hydride combinations (i.e. KHKHCO<sub>3</sub> units plus other positive fragments). The same compound was seen previously in a sample isolated from a  $K_2CO_3$  electrolytic cell [9].

The comparison of the positive ToF-SIMS spectrum of the KHCO<sub>3</sub> control with the KH KHCO<sub>3</sub> sample shown in Figs. 4 to 5 and 6, 7, respectively, demonstrates that the <sup>39</sup>K+ peak of the KH KHCO<sub>3</sub> sample may saturate the detector and give rise to a peak that is atypical of the natural abundance of <sup>41</sup>K. The natural abundance of <sup>41</sup>K is 6.7%; whereas, the observed <sup>41</sup>K abundance from the KHKHCO<sub>3</sub> sample is 53%. The high resolution mass assignment of the m/z = 41 peak of the KH KHCO<sub>3</sub> was consistent with <sup>41</sup>K, and no peak was observed at m/z = 42.98 ruling out  $H_2^{+}$ . Moreover, the natural abundance of <sup>41</sup>K was observed at the positive ToF-SIMS spectra of KHCO<sub>3</sub>, KhO<sub>3</sub>, and KI standards that were obtained with an ion current such that the <sup>39</sup>K peak intensity was an order of magnitude higher than that given for the KH KHCO<sub>3</sub> sample. The saturation of the <sup>39</sup>K peak of the positive ToF-SIMS spectrum by the KH KHCO<sub>3</sub> sample is indicative of a unique crystalline matrix [10].

The negative of-SIMS spectrum (m/e = 0-100) of the KHCO<sub>3</sub> (99.99%) sample and the KH KHCO<sub>3</sub> sample are shown in Fig. 8 and 9, respectively. The negative ion ToKSIM of the KH KHCO<sub>3</sub> sample was dominated by H<sub>2</sub>D<sub>7</sub>, and OH<sup>-</sup> peaks. A series of nonhydride containing negative ions {KCO<sub>3</sub>[K<sub>2</sub>CO<sub>3</sub>]<sub>n</sub><sup>-</sup> m/z = (99 + 138n)} was also present which implies that H<sub>2</sub> was eliminated to KH KHCO<sub>3</sub> during fragmentation of the compound KH KHCO<sub>3</sub>. Comparing the H<sup>-</sup> to O<sup>-</sup> ratio of the KH KHCO<sub>3</sub> sample to that of the KHCO<sub>3</sub> control sample, the H<sup>-</sup> peak was about an order of magnitude higher in the KH KHCO<sub>3</sub> sample.

## 3.1.3. ToF-SIMS of rubidium fluoro hydride sample

The positive ToF-SIMS spectrum (m/e = 0-100) of RbHF, the positive ToF-SIMS spectrum (m/e = 0-100) of RbF, the negative ToF-SIMS spectrum (m/e = 0-100) of RbHF, and the negative ToF-SIMS spectrum (m/e = 0-100) of RbF are shown in Figs. 10a-d, respectively. The positive ion spectrum of RbHF and that of the RbF control were dominated by the Rb<sup>+</sup> ion. Ga<sup>+</sup> m/z = 69,  $Rb_2^+$  m/z = 170, RbF<sup>+</sup> m/z = 104, and a series of positive ions Rb[RbF]<sub>n</sub> m/z = (85 + 104n) were also observed. The negative ion ToF-SIMS of RbHF was dominated by H<sup>-</sup> with a smaller F<sup>-</sup> peak. Fluorine alone dominated the negative ion ToF-SIMS of the RbF control. For both, O<sup>-</sup> m/z = 16, OH<sup>-</sup> m/z = 17, Cl<sup>-</sup> m/z = 35, RbF<sup>-</sup> m/z = 104, a series of negative ions  $F[RbF]_n^-$  m/z = (19 + 104n) were also observed.

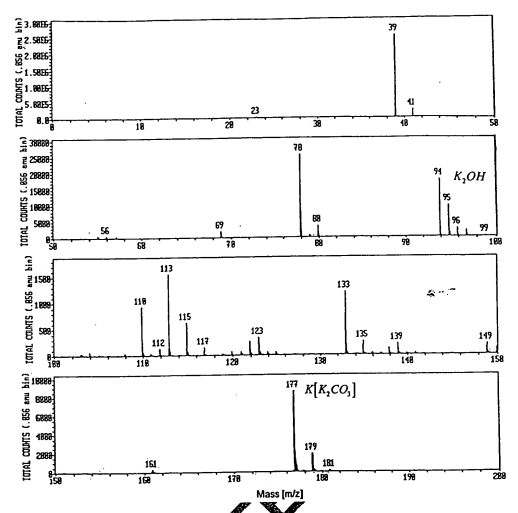


Fig. 4. The positive ToF-SIMS spectrum (m 200) of KHCO<sub>3</sub> (99.99%) where HC = hydrocarbon.

3.1.4. ToF-SIMS of strontium fluoro hydride same

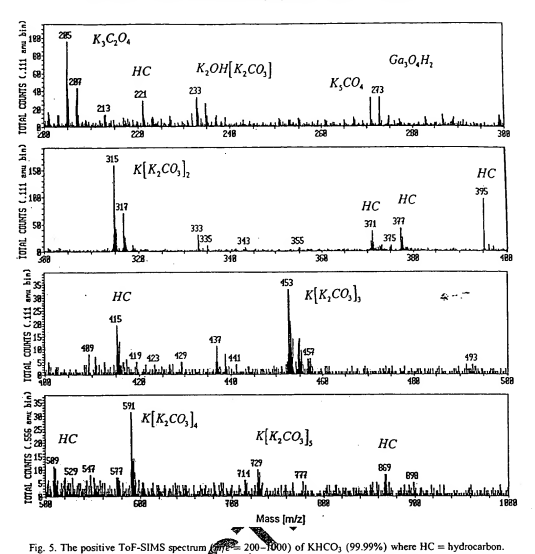
The positive ToF-SIMS spectrum obtained from the SrHF sample is shown in Fig. 11. The positive spectrum was dominated by the strontium peaks  $m_z = 88$  and a  $^{88}$ SrH+ m/z = 89 peak. Lithium sodiffication potassium, small hydrocarbon fragments such a  $C_2H_3$  m/z = 27 and  $C_2H_3$  m/z = 29,  $^{48}$ Ti+ m/z = 49,  $Ti_xO_y^+$ ,  $Ti_xO_yH^+$ ,  $K_2F^+$  m/z = 97  $^{88}$ SrOH+ m/z = 105,  $^{88}$ SrF+ m/z = 107 were also observed.

The positive spectrum of the  $SrF_2$  control was also dominated by the by the months of the smaller  $SrF_2$  control was also dominated by the  $SrF_2$  in the months  $SrF_3$  in  $SrF_4$  in

The negative ion ToF-SIMS of SrHF shown in Fig. 12 was dominated by  $H^-$  and  $F^-$  m/z = 19 of equal inten-

sity. Fluoride alone dominated the negative ion ToF-SIMS of the SrF<sub>2</sub> control. For both samples, smaller  $O^- m/z = 16$ ,  $OH^- m/z = 17$ , hydrocarbon fragment peaks such as  $C^- m/z = 12$  and  $CH^- m/z = 13$ ,  $CI^- m/z = 35$ ,  $SrF_2^- m/z = 126$ , and  $I^- m/z = 127$  were observed. A hydride peak which was significantly smaller than the  $O^- m/z = 16$  peak was observed in the control.  $Ti_xO_yH^-$  was observed in the SrHF sample. The hydrocarbon peaks were much more intense in the control.

The negative ToF-SIMS relative sensitivity factors (RSF) for the halides are all about equivalent. The hydrino hydride ion is in the same group as the halide ions. Thus, its RSF is projected to be equivalent to that of the halides. Therefore, the atomic percentage of hydrino hydride ion may be determined by comparison of its intensity with that of the fluoride ion of the product SrHF. The atomic percentage of hydrino hydride ion is equivalent to that of fluoride as shown in the negative ToF-SIMS. Since essentially no other cations are present, the positive and negative ToF-SIMS



indicates that SrHF is the product of the gas call reaction of SrF2 in the presence of atomic hydrogen and strong in metal catalyst.

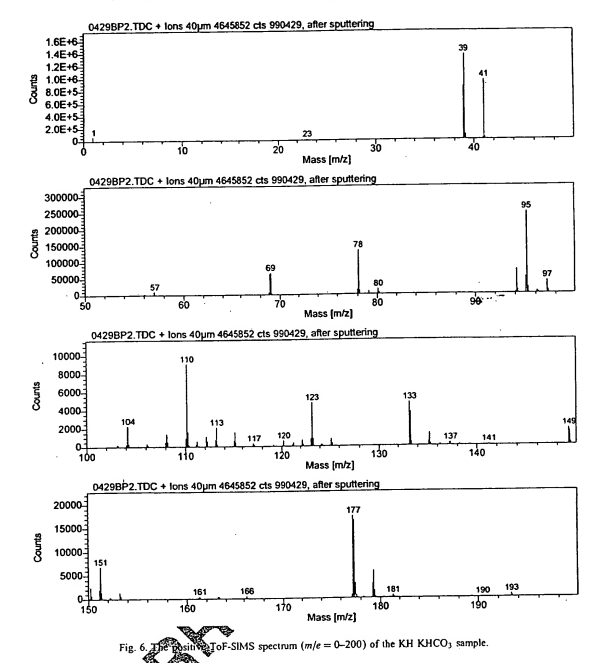
## 3.2. XPS

3.2.1. XPS of potassium iodo hydride sample
A survey spectrum was obtained over the region  $E_b = 0$ to 1200 eV. The primase element peaks allowed for the determination of all of the elements present in MHX and the control Ma The survey spectrum also detected shifts in the binding energies of the elements which had implications to the identity of the compound containing the elements.

The XPS survey scan of KI and KHI are shown in Figs. 13a and b, respectively. C1s at 284.5 eV was used as the internal standard for KHI and the control KI. The major species present in the KHI sample and the control are potassium and iodide. Trace small amounts of carbonate carbon and oxygen were also identified in the KHI sample.

The 0-100 eV binding energy region of a high resolution XPS spectrum of KI and KHI are shown in Figs. 14a and b, respectively. Peaks centered at 21 and 37 eV which do not correspond to any other primary element peaks were observed in the case of the KHI sample. The intensity and shift match shifted K3 s and K3 p. Hydrogen is the only element which does not have primary element peaks; thus, it is the only candidate to produce the shifted peaks. These peaks may be shifted by a highly binding hydride ion H<sup>-</sup>(1/6) with a binding energy of 22.8 eV given by Eq. (A.12) that bonds to potassium K3 p and shifts the peak to this energy. In this case, the K3 s is similarly shifted.

## HE 1158



3.2.2. XPS of potassium chioro liveride sample

A survey spectrum was obtained over the region  $E_b = 0$  to 1200 eV. The primary element peaks allowed for the determination of all of the elements present in the magenta crystals of the KHC sample and the control KCl. The survey spectrum also detected shifts in the binding energies of the elements which had implications to the identity of the compound containing the elements.

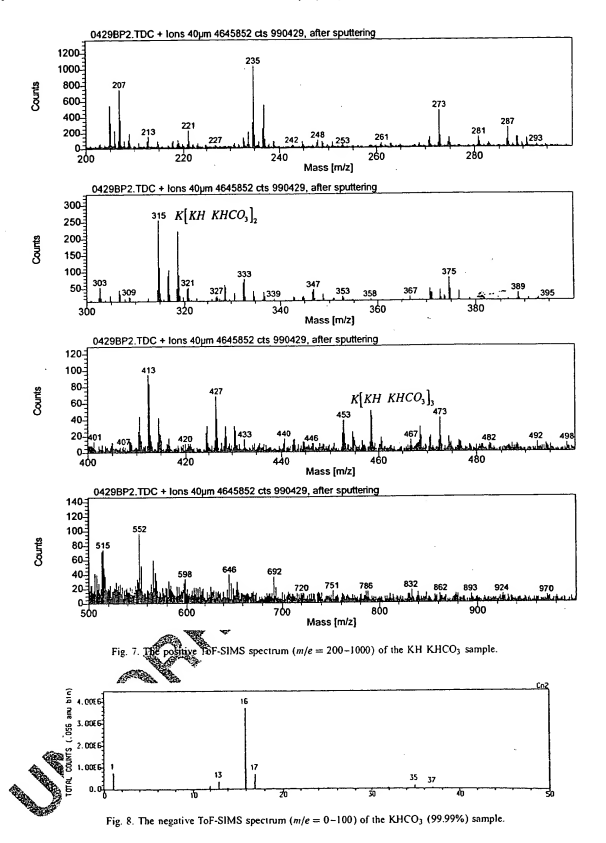
The XPS survey scan of the KHCl sample and the KCl control sample are shown in Figs. 15 and 16, respectively. Cls at 284.6 eV was used as the internal standard for the

KHCl sample and the control KCl sample. The major species observed in the KHCl sample and the control were potassium and chlorine. Trace small amounts of carbon, oxygen, fluorine, iodine, and silicon were also identified in the KHCl sample. The identifying peaks of the primary elements and their binding energies are: FKL23L23 at 831.0 eV, F1s at 688.4 eV and at 682.8 eV, O1s at 530.6 eV, K2s at 377.2 eV, K2p1/2 at 295.4 eV, K2p3/2 at 292.5 V, C1s at 284.6 eV, C12s at 268.9 eV, C12p1/2 at 199.5 eV, C12p3/2 at 198.0 eV, Si2s at 152.4 eV, and Si2p3/2 at 101.9 eV. The K 3p and K 3s of the KHCl sample occurred at 16.7 and 32.8 eV,





R.L. Mills et al. | International Journal of Hydrogen Energy 000 (2000) 000-000



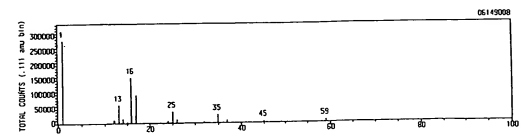


Fig. 9. The negative ToF-SIMS spectrum (m/e = 0-100) of the KH KHCO<sub>3</sub> sample.

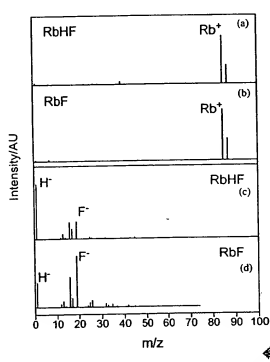


Fig. 10. (a) The positive ToF-SIMS spectrum (m/e = 0) of RbHF. (b) The positive ToF-SIMS spectrum (m/e=0-10) of Rb (c) The negative ToF-SIMS spectrum (m/e = 0 - 100) de RbH (d) The negative ToF-SIMS spectrum (m/e = 0<sub>2</sub>

ntrol KCl occurred respectively. The K3p and K3 s of the

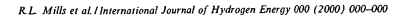
at 17.0 and 32.8 eV, respectively.

No elements were present in the survey scan which could be assigned to peaks in the low binding energy region with the exception of the K 3h and 33s peaks at 16.7 and 32.8 eV, respectively, and the 14de, and 14ds/2 peaks at 51.8 and 49.7 eV respectively. Accordingly, any other peaks in this region must be due to nevel species. The 0-125 eV binding energy region of a high resolution XPS spectrum of the KHCl sample and the control KCl sample are shown in Figs. 17 and 18, respectively. The XPS spectrum of the KHE sample differs from that of KCl by having an additional feature at 36.7 eV. The XPS peak centered at 36.7 eV that does not correspond to any other primary element peak

may correspond to the  $H^-(n = 1/8)E_b = 36.1$  eV hydride ion given by Eq. (A.12) where  $E_b$  is the predicted vacuum binding energy. Also the K3s and K3 p peaks in the KHCl sample are wider when compared to the K3 s and K3 p of the KCl control. The Cl3p at 16 eV is merged with the K3 p of the KHCl sample, but the peaks are separated in the control taken at the same resolution. This clearly indicates that the environment of K in the KHC sample is different from that of KCI. The data further male at the formation of a novel compound.

3.2.3. XPS of alkali halido hydrides and alkaline earth

halido hydrides samples hinding energies and features of core level electrons of a series of alkali and alkaline earth halido hydrides (KHF, KHCI, KHBA KHI, RBHF, RbHCI, RbHBr, RbHI, CsHF, CsHCI, CsHB, CsHCI, CaHCI, CaHCI, CaHBr, CaHI, SrHF, SrHCI, and SrHBr) were analyzed by XPS. The local structure of metal halides and metal halido hydrides was investigated by studying the metal core levels including K 2p, Rb 3d, Cs de Caro, and Sr 3d, and halogen core level including F , C\$2p Br 2p, and I 3d. As atomic hydrogen undergoes eaction with a catalyst to form a lower-energy hydrogen species which subsequently reacts with the metal center in a halide compound, alterations in the electronic structure of the metal such as changes in core level binding energies and spin-orbital energies relative to the starting halide are expected. In order to compare the full width at half maximum (FWHM) of the peaks, the difference between spin-orbit splitting in the core level was determined using curve fitting in the same or a close energy range. In some cases, the absolute core level binding energy was not calibrated, which did not affect the validity of the comparison of FWHM values. The results of the determination of the binding energies of selected core level electrons, full width at half maximum of the peaks, and energy of spin-orbital splitting for alkali halido hydrides, calcium halido hydrides, and strontium halido hydrides compared with the corresponding alkali halides, calcium halides, and strontium halides are listed in Tables 1-3, respectively. The XPS spectra of the K 2p core level in KI, KHI appear in Figs. 19a and b, respectively. The XPS spectra of the 1 3d core level in K1 and KHI appear in Figs. 20a and b, respectively. The XPS spectra of the Ca 2p core level in CaBr2 and CaHBr appear in Figs. 21a and



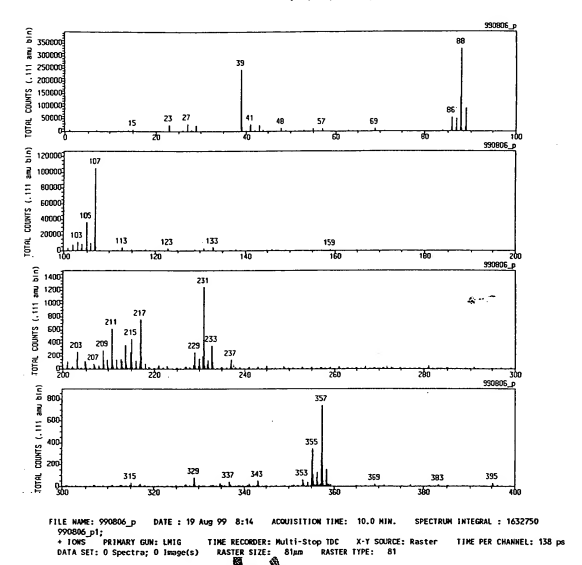


Fig. 11. The positive Top SIMS spectrum obtained from the SrHF sample.

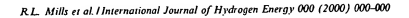
b, respectively. The XPS spectra of the Symptote level in  $SrBr_2$  and SrHBr appear in Figs. 22a and b, respectively. The XPS spectra of the Br 2p core level in  $SrBr_2$  and SrHBr appear in Figs. 23a and b, respectively.

It is clear that the FWIM of the metal core level peaks in alkali and alkaline earth halido hydrides is broader than that in the corresponding latities. The magnitude of broadening ranges from 0.3 to 0.9 eV, depending on compound. The trend for the broadening effect follows the sequence: iodo hydrides brome hydrides > fluoro hydrides chloro hydrides line ontrast, the halogen core level FWHM in both alkali and alkaline earth halido hydrides is broadened by only about 0.2 to 0.3 eV compared to the corresponding halide. Comparing the alterations in the metal core levels versus the halogen core level indicates that the lower-energy

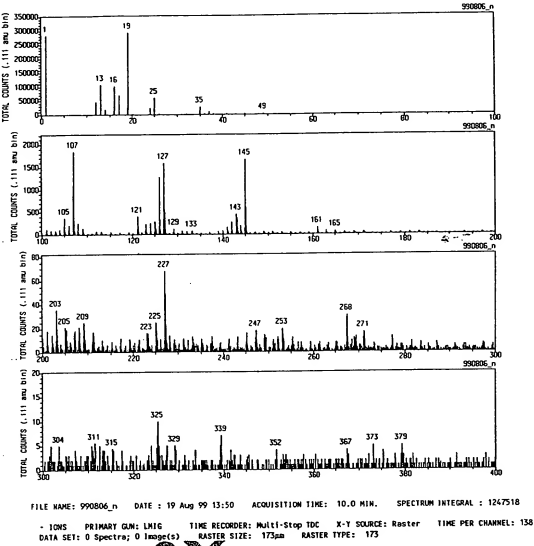
hydrogen species is bound to the metal center of the alkali or alkaline earth halide. This binding influences the metal core level with little perturbation of the halogen core level.

Each of the spectra of potassium iodo hydride, calcium bromo hydride, and strontium bromo hydride were curve fit with one spin-orbit splitting component having a similar FWHM and energy separation as that of the starting material potassium iodide, calcium bromide, and strontium bromide, respectively. An additional spin-orbit splitting component had to be added to each of potassium iodo hydride, calcium bromo hydride, and strontium bromo hydride in order to obtain a good curve fit of the K 2p, Ca 2p, and Sr 3d spectra. In each case, the second component of spin-orbit splitting is assigned to the formation of the alkali or





# AMALYTICAL SERVICES GROUP SURFACE AMALYSIS LABORATORY 1:609-490-1090 F:609-490-1066



roF-SIMS spectrum obtained from the SrHF sample.

alkaline earth metal halido hydride KHI, CaHBr. and SrHBr, respectively. The present of the novel hydride ion shifts the K 2p, Ca 2p and Sr3d peaks to lower binding energies relative to the corresponding peaks of KI, CaBr<sub>2</sub>, and SrBr<sub>2</sub>, respectively.

The XPS data clearly indicates a change in the electronic structure at the metal core level and different bonding in the metal halfdre hydrodes relative to that in the corresponding metal halides it strongly suggests the formation of a novel metal hydride which is consistent with the supporting data provided by XPS given above and NMR, ToF-SIMS, and gas chromatography/mass spectroscopy given in the respective sections.

## 3.3. NMR

To eliminate the possibility that the alkali halide MX influenced the local environment of the ordinary alkali hydride MH to produce an NMR resonance that was shifted upfield relative to MH alone, controls comprising MH and a MH/MX mixture were run.

## 3.3.1. NMR of potassium iodo hydride sample

The <sup>1</sup>H MAS NMR spectra of the KHI sample, the control comprising an equal molar mixture of KH and Kl, and control KH relative to external tetramethylsilane (TMS) are shown in Figs. 24a, b, and c, respectively. Ordinary hy-



R.L. Mills et al. | International Journal of Hydrogen Energy 000 (2000) 000-000

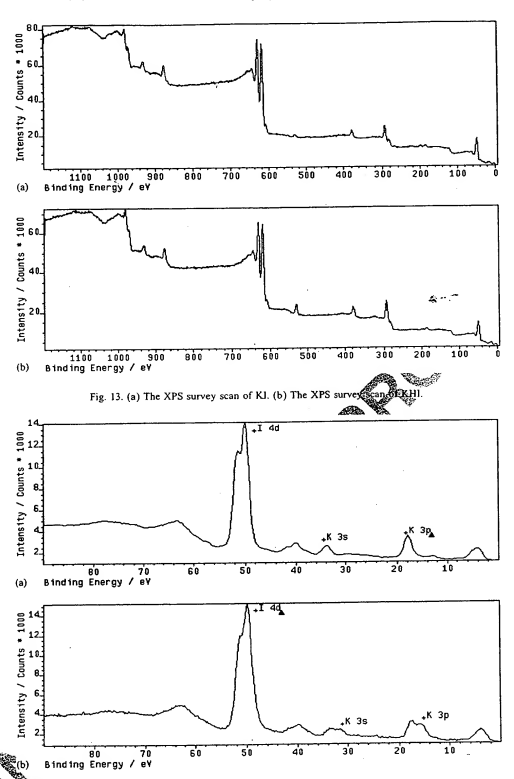


Fig. 14. (a) The 0-100 eV binding energy region XPS spectrum of KI. (b) The 0-100 eV binding energy region XPS spectrum of KHI.

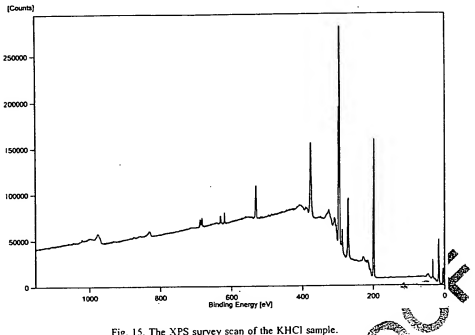


Fig. 15. The XPS survey scan of the KHCl sample.

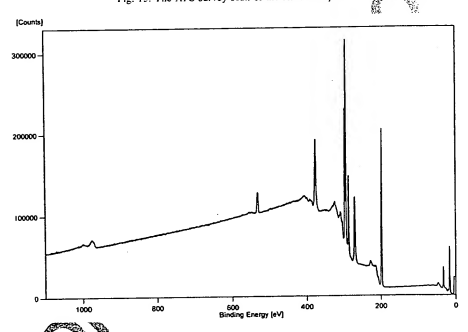


Fig. 16. The XPS survey scan of the KCl control sample.

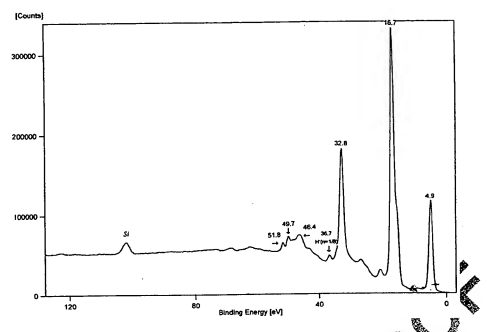
dride ion has a resonance at 1.1 and 0.8 ppm in the KH/KI mixture and in KH alone as shown in Figs. 24b and c, respectively the additional peak at 4.5-4.6 ppm is assigned to KOH tormed from air exposure of KH during sample handling. The spin speed was varied to confirm real peaks versus side bands. The latter changed position with spin speed, the former were independent of spin speed. The unlabeled peaks shown in Figure 24a were found to be side-

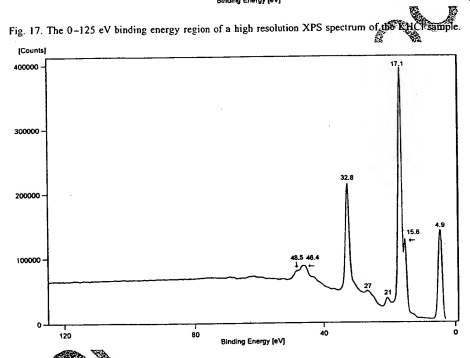
The presence of KI does not shift the resonance of ordinary hydride. The resonance at 0.9 ppm which is assigned to ordinary hydride ion was observed in the spectrum of the KHI sample as shown in Fig. 24a. The distinct 0.8 and 1.1 ppm resonances could not be resolved if they were present.





## R.L. Mills et al. | International Journal of Hydrogen Energy 000 (2000) 000-000





V binding energy region of a high resolution XPS spectrum of the KCl control sample. Fig. 18. Th

A large distinct unfield resonance was observed at -3.2 ppm which was not observed in either control. This upfield shifted peak is consistent with a hydride ion with a smaller radius as compared with ordinary hydride since a smaller radius increases the shielding or diamagnetism. The  $-3.2\,$  ppm peak is assigned to a novel hydride ion that has a smaller radius than that of ordinary hydride ion since the shift was extraordinarily far upfield in the case of the KHI sample.

## 3.3.2. NMR of potassium chloro hydride sample

The <sup>1</sup>H MAS NMR spectra of the KHCl sample, the control comprising an equal molar mixture of KH and KCl, and the control KH relative to external tetramethylsilane (TMS) are shown in Figs. 25a, b, and c, respectively. Ordinary hydride ion has a resonance at 1.1 and 0.8 ppm in the KH/KCl mixture and in KH alone as shown in Figs. 25b and c, respectively. The additional peak at 6 ppm is assigned

## HE 1158

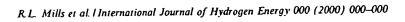


Table 1
The results of the determination of the binding energies of selected core level electrons, full width at half maximum of the peaks, and energy of spin-orbit splitting for alkali halido hydrides compared with the corresponding alkali halides

Compound	Peak	Binding energy (eV) <sup>a</sup>	Full width at half maximum (FWHM) (eV)	Energy of spin- orbit splitting (eV
KF	K 2p3/2	298.37	2.24	2.80
	K 2p1/2	301.11	2.27	
	Fls	688.97	2.33	
КНБ	K 2p3/2	296.12	2.61	2.72
	K 2p1/2	298.84	2.61	
	F 1s	687.60	2.20	
KCI	K 2p3/2	296.87	1.88	2.75
	K 2p1/2	299.62	1.97	•
	Cl 2p3/2	202.27	1.82	1.68
	Cl 2p1/2	203.95	1.58	A
кнсі	K 2p3/2	297.09	1.91	<b>62.7</b>
Mici	K 2p1/2	299.85	1.98	
	Cl 2p3/2	202.57	1.77	.62
	Cl 2p1/2	204.19	1.60	
KBr	K 2p3/2	297.15	1.63	2.75
KDI	K 2p3/2 K 2p1/2	299.90	1.72	
	Br 3p3/2	186.34	2.55	6.87
	Br 3p1/2	193.07	2.39	
КНВг	K 2p3/2	296.90	2.10	2.76
Kibi	K 2p1/2	299.66	1.86	
	Br 3p3/2	186.18		6.79
	Br 3p1/2	192.87		
кі	K 2p <sub>3/2</sub>	294.44	1.84	2.75
	K 2p <sub>1/2</sub>	297.19	· 🕦.	
	1 3d <sub>3/2</sub>	620.37	2.07	11.53
	1 3d <sub>3/2</sub>	631.90	2.14	
кні	K 2p <sub>3/2</sub>	294.37	1.83	2.77
Na.i	K 2p <sub>1/2</sub>	29.14	1.81	
	K 2p <sub>3/2</sub>	2921	1.87	
	K 2p <sub>1/2</sub>	2295	1.78	3.00
	1 3d <sub>5/2</sub>	621.08	2.17	11.52
	1 3d <sub>3/2</sub>	632.60	2.20	
RbF	Rb 3d <sub>5/2</sub>	115.00	2.28	1.55
	R4F3d3/2	116.55	2.08	
•		688.35	2.30	
RbHF	Rh3dsn	114.65	1.95	1.55
	RB3d3/2	116.37	1.56	
	Rb 3d <sub>5/2</sub>	114.02	2.24	- 1.56
	Rb 3d <sub>3/2</sub>	115.58	1.88	1.50
	Fls	688.35	2.10	
RbCla A	Rb 3d <sub>5/2</sub>	115.17	1.84	1.55
	Rb 3d <sub>3/2</sub>	116.72	1.77	



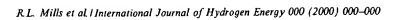
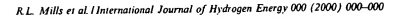


Table 1. (Continued)

Compound	Peak	Binding energy (eV) <sup>a</sup>	Full width at half maximum (FWHM) (eV)	Energy of spin- orbit splitting (eV
	Cl 2p <sub>3/2</sub>	203.39	1.75	1.66
	Cl 2p <sub>1/2</sub>	205.05	1.69	
RЬНСІ	Rb 3d <sub>5/2</sub>	114.30	2.17	1.56
	Rb 3d <sub>3/2</sub>	115.86	2.02	
	Cl 2p <sub>3/2</sub>	202.59	2.01	1.72
	Cl 2p <sub>1/2</sub>	204.33	1.70	
RbBr	Rb 3d <sub>5/2</sub>	114.46	1.76	1.51
KUDI	Rb 3d <sub>3/2</sub>	115.97	1.83	
	Br 3p <sub>3/2</sub>	186.56	2.88	6.66
	Br 3p <sub>1/2</sub>	193.22	1.75	0.00
DLUD-	Rb 3d <sub>5/2</sub>	114.54	1.74	1.56
RbНВг	Rb 3d <sub>3/2</sub>	116.0	1.85	1.50
	Br 3p <sub>3/2</sub>	186.65	2.70	. 63
	ът эр <sub>3/2</sub> Вт 3р <sub>1/2</sub>	193.38	1.69	
DLI	Rb 3d <sub>5/2</sub>	114.54	1.99	
RbI	Rb 3d <sub>3/2</sub>	116.09	1.81	
		623.72	2.04	11.49
	1 3d <sub>5/2</sub> 1 3d <sub>3/2</sub>	635.21	2.07	11.42
			2.31	1.56
RЬНІ	Rb 3d <sub>5/2</sub> Rb 3d <sub>3/2</sub>	114.29 115.85	1.89	1.50
				11.50
	I 3d <sub>5/2</sub> I 3d <sub>3/2</sub>	623.28 634.78	2.10 2.17	11.50
C-E			2.3	
Cs <b>F</b>	Cs 3d <sub>5/2</sub>	746.18		
	F ls	690.55		
CsHF	Cs 3d <sub>5/2</sub>	741.30	192.2	
	Cs 3d <sub>5/2</sub>	738.63		
	F Is	687.81	3.01	
	F 1s	684.23	2.60	
CsC1	Cs 3d <sub>5/2</sub>	7471	2.04	
	Cl 2p <sub>3/2</sub>	205	1.65	1.50
	Cl 2p <sub>1/2</sub>	200218	1.43	
CsHCl	Cs 3d <sub>5/2</sub>	J7437	2.35	
	Cl 2p <sub>3/2</sub>	203.60	1.73	1.56
	Cl 2p <sub>1/2</sub>	205.24	1.58	
CsBr	Cs 3d <sub>3/2</sub>	745.01	1.78	
	Br. Dn.	188.33	2.35	6.68
	Br2p <sub>1/2</sub>	195.01	2.13	
Coup.		728.79	2.40	
CsHBr	Cs 3d <sub>5/2</sub>	140.17	2.70	*
	The state of the s	186.38	2.45	6.63
	Br 2p <sub>3/2</sub> Br 2p <sub>1/2</sub>	193.01	2.08	0.05
*Uncalibrated, o		175.01	2.00	

## HE 1158



The results of the determination of binding energies of selected core level electrons, full width at half maximum of the peaks, and energy of spin-orbit splitting for calcium halido hydrides compared with the corresponding calcium halides

Compound	Peak	Binding energy (eV) <sup>a</sup>	Full width at Half Maximum (FWHM) (eV)	Energy of spin- orbit splitting (eV)
CaCl <sub>2</sub>	Ca 2p <sub>3/2</sub>	352.18	2.25	3.51
	Ca 2p <sub>1/2</sub>	355.69	2.19	
	Cl 2p <sub>3/2</sub>	203.12	2.28	1.81
	Cl 2p <sub>1/2</sub>	204.93	1.86	
CaHCl	Ca 2p <sub>3/2</sub>	351.95	2.88	3.67
	Ca 2p <sub>1/2</sub>	355.62	2.34	
	C1 2p <sub>3/2</sub>	202.75	2.25	1.57
	Cl 2p <sub>1/2</sub>	204.32	2.01	
CaBr <sub>2</sub>	Ca 2p <sub>3/2</sub>	348.24	1.79	351
•	Ca 2p <sub>1/2</sub>	351.75	1.83	
	Вг 3р <sub>3/2</sub>	182.86	2.62	6.88
	Br 3p <sub>1/2</sub>	189.54	2.36	
СаНВг	Ca 2p <sub>3/2</sub>	347.36	2.38	3.59
	Ca 2p <sub>1/2</sub>	350.95	2.02	
	Ca 2p <sub>3/2</sub>	349.06	1.85	3.41
	Ca 2p <sub>1/2</sub>	352.47	1.60	
	Br 3p <sub>3/2</sub>	182.86	3.16	6.66
	Br 3p <sub>1/2</sub>	189.52	2.97	
Cal <sub>2</sub>	Ca 2p <sub>3/2</sub>	356.08	215	2.54
	Ca 2p <sub>1/2</sub>	358.62	2.13	
	1 3d <sub>5/2</sub>	624.27	2.28	11.51
	1 3d <sub>3/2</sub>	635.78	29.33	
CaHI	Ca 2p <sub>3/2</sub>	352.05	3.08	3.86
	Ca 2p <sub>1/2</sub>	355.91	2.63	
	1 3d <sub>5/2</sub>	4.0	2.57	11.54
	1 3d <sub>3/2</sub>	63.57	2.46	

\*Uncalibrated, except for CaBr2 and

to KHCO<sub>3</sub> formed from all exposure of K during sample handling. The additional sharp peak at 4.3 ppm shown in Fig. 25b is assigned to water in the KCl crystals. The ad-

ditional broad peak at 60 ppm shown in Fig. 25c is assigned KOH formed from air exposure of KH during sample handling

The presence of KCl does not shift the resonance of ordinary hydride. The resonance at 1.1 ppm which is assigned to ordinary hydride ion was observed in the spectrum of the KHCP sample as shown in Fig. 25a. The distinct 0.8 and 1.1 ppm resonances could not be resolved if they were

present. A large distinct upfield resonance was observed at -4.6 ppm which was not observed in either control. This upfield shifted peak is consistent with a hydride ion with a smaller radius as compared with ordinary hydride since a smaller radius increases the shielding or diamagnetism. The -4.6 ppm peak is assigned to a novel hydride ion that has a smaller radius than that of ordinary hydride ion since the shift was extraordinarily far upfield in the case of the KHCl sample. An additional water peak was found is other samples in which the novel hydride peak was observed. The presence of a water peak may demonstrate that the novel

Table 3 The results of the determination of the binding energies of selected core level electrons, full width at half maximum of the peaks, and energy of spin-orbit splitting for strontium halido hydrides compared with the corresponding strontium halides

Compound	Peak	Binding energy (eV) <sup>a</sup>	Full width at half maximum (FWHM) (eV)	Energy of spin- orbit splitting (eV)
SrF <sub>2</sub>	Sr 3d <sub>5/2</sub>	140.46	2.13	1.81
	Sr 3d <sub>3/2</sub>	142.27	1.77	
	Fls	691.12	2.11	
SrH <b>F</b>	Sr 3d <sub>5/2</sub>	139.20	2.48	1.91
	Sr 3d <sub>3/2</sub>	141.11	1.95	
	F 1s	689.88	2.42	
SrCl₂	Sr 3d <sub>5/2</sub>	139.45	2.36	1.86
	Sr 3d <sub>3/2</sub>	141.31	1.88	
	Cl 2p <sub>3/2</sub>	203.61	2.06	7.70
	Cl 2p <sub>1/2</sub>	205.31	1.95	
SrHCl	Sr 3d <sub>5/2</sub>	138.71	2.61	1.81
	Sr 3d <sub>3/2</sub>	140.52	1.99	
	Cl 2p <sub>3/2</sub>	202.98	2.06	1.72
•	Cl 2p <sub>1/2</sub>	204.70	1.89	•
SrBr <sub>2</sub>	Sr 3d <sub>5/2</sub>	139.52	1.58	1.7
	Sr 3d <sub>3/2</sub>	141.22	1.65	
	Br 3p <sub>3/2</sub>	187.45	2.50	6.77
	Br 3p <sub>1/2</sub>	194.22	2	
SrHBr	Sr 3d <sub>5/2</sub>	138.56		1.96
	Sr 3d <sub>3/2</sub>	140.52	4.62	
	Sr 3d <sub>5/2</sub>	136.69	95	2.60
	Sr 3d <sub>3/2</sub>	139.29	1.55	
	Br 3p <sub>3/2</sub>	186.56	2.77	6.68
	Br 3p <sub>1/2</sub>	19324	2.43	

<sup>&</sup>lt;sup>8</sup>Uncalibrated, except for SrBr<sub>2</sub> and SrHB

hydride is stable to water, in the case that the water is not associated with unreacted KCl atine. This could partially explain why the KHCl sample was observed to be insoluble in

3.3.3. NMR of parassium thromo hydride sample

The <sup>1</sup>H MAS NMR spectra of the KHBr sample, the control comprising an equal molar mixture of KH and KBr, and the control KH relative to external tetramethylsilane (TMS) are shown in Figs. 26a, b, and c, respectively. Ordinary hydride ion has a resonance at 1.1 and 0.8 ppm in the KH/KBr mixture and in KH alone as shown in Figs. 26b and 26c, respectively. The additional sharp peaks at 4.3 and 5.9 ppm shown in Fig. 26a are assigned to water and KHCO3 formed

from air exposure of K during sample handling. The additional sharp peak at 4.2 ppm shown in Fig. 26b is assigned to water in the KBr crystals. The additional broad peak at 4.6 ppm shown in Fig. 26c is assigned KOH formed from air exposure of KH during sample handling.

The presence of KBr does not shift the resonance of ordinary hydride. The resonance at 1.2 ppm which is assigned to ordinary hydride ion was observed in the spectrum of the KHBr sample as shown in Fig. 26a. The distinct 0.8 and 1.1 ppm resonances could not be resolved if they were present. A large distinct upfield resonance was observed at -4.1 ppm which was not observed in either control. This upfield shifted peak is consistent with a hydride ion with a smaller radius as compared with ordinary hydride since a

## HE 1158

R.L. Mills et al. I International Journal of Hydrogen Energy 000 (2000) 000-000

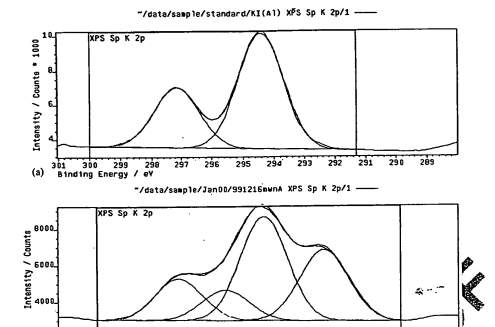


Fig. 19. (a) The XPS spectra of the K 2p core level in KI. (b) The XPS spectra of the K 2p core level in KHI.

295

294

293

292

296

297

301 300 299 298 Binding Energy / eY

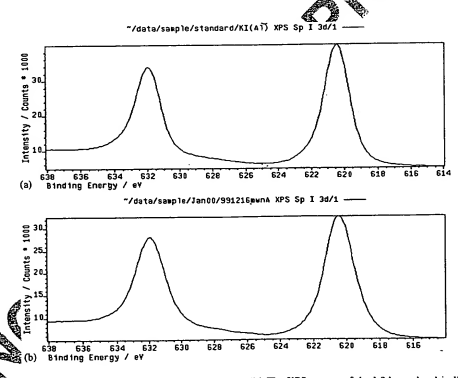


Fig. 20. (a) The XPS spectra of the I 3d core level in KI. (b) The XPS spectra of the I 3d core level in KHI.



9

## R.L. Mills et al. | International Journal of Hydrogen Energy 000 (2000) 000-000

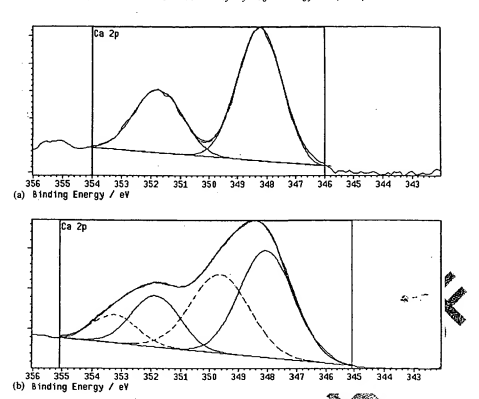


Fig. 21. (a) The XPS spectra of the Ca 2p core level in CaBr<sub>2</sub>. (b) The XPS spectra of the Ca 2p core level in CaHBr.

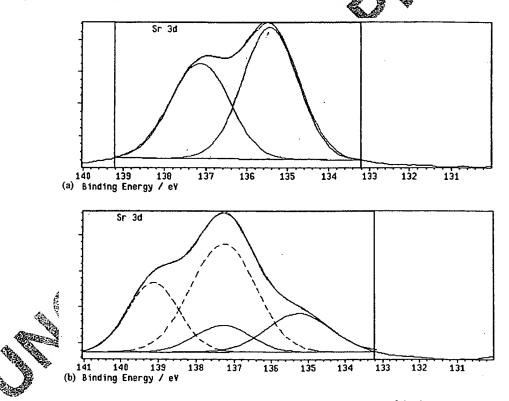
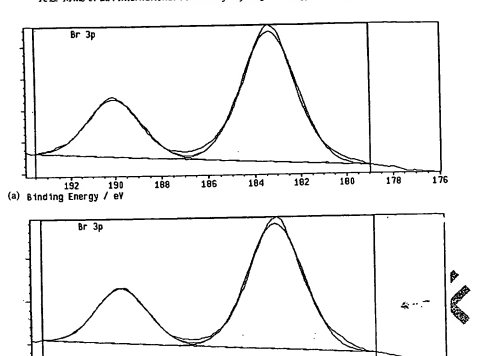


Fig. 22. (a) The XPS spectra of the Sr 3d core level in SrBr<sub>2</sub>. (b) The XPS spectra of the Sr 3d core level in SrHBr.



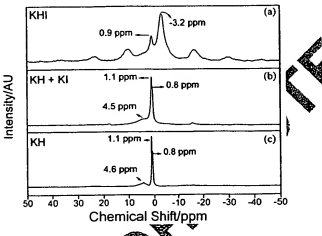




of the Br 2p core level in SrHBr. Fig. 23. (a) The XPS spectra of the Br 2p core level in SrBr2. (b) The XPS spectra

186

184



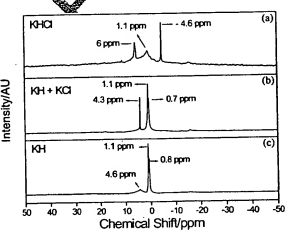
192 (b) Binding Energy / eV

190

188

Fig. 24. (a) The <sup>1</sup>H MAS NMR spectrum of KHI relative to external tetramethylsilane (TMS). (b) The IB MAS NMR spectrum of the control comprising an equal motor mixture of KH and KI relative to external tetramethylsilane (EMS). (c) The <sup>1</sup>H MAS NMR spectrum of the control KH relative to external tetramethylsilane (TMS).

smaller addits increases the shielding or diamagnetism. The -All ppin peak is assigned to a novel hydride ion that has a smaller radius than that of ordinary hydride ion since the shift was extraordinarily far upfield in the case of the KHBr sample.



180

182

Fig. 25. (a) The <sup>1</sup>H MAS NMR spectrum of KHCl relative to external tetramethylsilane (TMS). (b) The <sup>1</sup>H MAS NMR spectrum of the control comprising an equal molar mixture of KH and KCl relative to external tetramethylsilane (TMS). (c) The <sup>1</sup>H MAS NMR spectrum of the control KH relative to external tetramethylsilane (TMS).

## 3.3.4. NMR of rubidium fluoro hydride sample

The 1H MAS NMR spectrum of RbHF relative to external tetramethylsilane (TMS) is shown in Fig. 27a. Distinguishable resonances were observed at 1.2 and -4.4 ppm. The

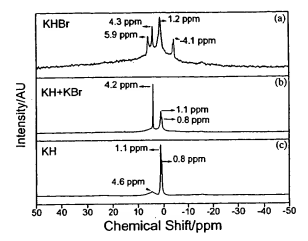


Fig. 26. (a) The <sup>1</sup>H MAS NMR spectrum of KHBr relative to external tetramethylsilane (TMS). (b) The <sup>1</sup>H MAS NMR spectrum of the control comprising an equal molar mixture of KH and KBr relative to external tetramethylsilane (TMS). (c) The <sup>1</sup>H MAS NMR spectrum of the control KH relative to external tetramethylsilane (TMS).

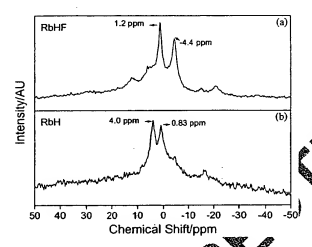


Fig. 27. (a) The <sup>1</sup>H MAS NMR spectrum of RbH relative to external tetramethylsilane (TMS). (b) The H MAS NMR spectrum of the control RbH relative to external tetramethylsilane (TMS).

upfield peak is assigned to a novel hydride ion of RbHF. The down field shifted peak may be ordinary hydride in a different chemical environment. The 1H MAS NMR spectrum of the control RbH relative to external tetramethylsilane (TMS) ris shown in Fig. 27b. The 0.83 ppm peak is assigned to ordinary hydride ion of RbH. The peak at 4.0 ppm is assigned to RbOH formed from air exposure of RbH during sample handling.

The upfield shifted peak observed in the RbHF sample is consistent with a hydride ion with a smaller radius as compared with ordinary hydride since a smaller radius increases the shielding or diamagnetism. The -4.4 ppm peak is assigned to a novel hydride ion that has a smaller radius than that of ordinary hydride ion since the shift was extraordinarily far upfield in the case of the RbHF sample.

#### 3.3.5. NMR of strontium bromo hydride sample

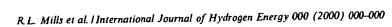
The SrHBr sample was not spun; so, the hydride peaks were not resolved. Instead valuable 'H-'H separation data as well as <sup>1</sup>H content was obtained. The <sup>1</sup>H MAS NMR spectra of SrHBr relative to external H2O is shown in Fig. 28. The sample contained a significant amount of hydrogen with an integration of 0.0983 mmoles corresponding to about 0.02 wt% <sup>1</sup>H in the sample. There are two components: a narrow component with a peak at 6.3 ppm having an area of approximately 1/4th of the total signal, and a broad signal with a width of about 60 kHz The broadening of the peak corresponds to  ${}^{1}H^{-1}H$  separation of under 1.5 Å. This is very significant given that the unit cell (orthorhombic  $P_{nma}$  form) parameters for ordinary stroptium hydride are a = 7.358 Å, b = 6.377 Å and c = 3.882 Å. The seven short M-H distances are 2.49 Å. The two long M-H distances are 3.06 Å. The Haradius 1.39 Å [11]. The data is consistent with a novel hydride ion of a greatly reduced radius.

## 3.4. Gas chromaingraphy (GC)

The gas chromatograph of the normal hydrogen gave the retention time for para hydrogen and ortho hydrogen as 14.5 and \$5.5 min, respectively. Control SrBr2 showed no hydrogen release upon heating to above 600°C. The gas chromatograph of the dihydrino (see Appendix) or hydrogen resed from the SrHBr sample when it was heated to above 600°C is shown in Fig. 29. This chromatograph is representative of the results of the determination of dihydrino or hydrogen released from each sample of MHX or MHMX when heated to 600°C. In the case of a very stable hydride, no hydrogen may be released, thus, the amount observed in the minimum content. Table 4 gives a results which are representative of the minimum dihydrino or hydrogen content of novel hydride compounds determined by gas chromatograph of the gas released from each sample when it was heated to above 600°C. Each control, MX or MX2, showed no hydrogen release upon heating to above 600°C.

## 3.5. Mass spectroscopy (MS)

The dihydrino (see Appendix) was identified in the gas released by thermal decomposition of solid samples by mass spectroscopy. Dihydrino was detected as a species with a mass to charge ratio of two (m/e = 2) that has a higher ionization potential than that of normal hydrogen by recording the ion current as a function of the electron gun energy. The intensity as a function of time for masses m/e = 1, m/e = 2,





**HE 1158** 

8000



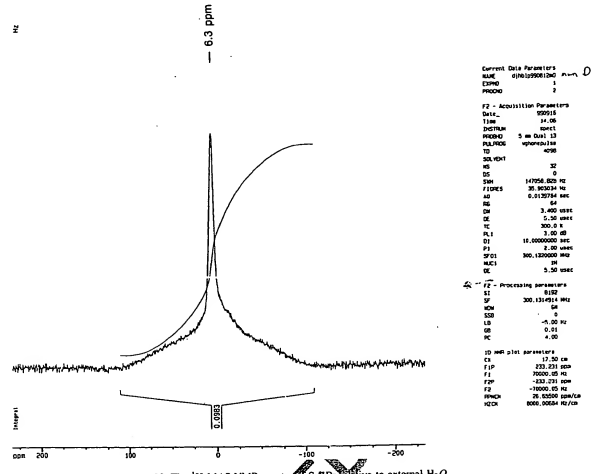


Fig. 28. The <sup>1</sup>H MAS NMR spectra of SchBr relative to external H<sub>2</sub>O.

and m/e = 3 obtained while changing the ionization potential (IP) of the mass spectrometer from 30 to 70 eV trapure hydrogen is shown in Fig. 30. Upon the easing the ionization potential from 30 to 70 eV, the ally the n/e = 2 ion current for the ultrapure hydrogen uncerted by a factor of less than 2.

The intensity as a function of time for masses m/e = 1, m/e = 2, and m/e = 3 obtained while changing the ionization potential (IP) of the mass spectrometer from 30 to 70 eV for gas released from thermal decomposition of the KHI sample is shown in Fig. 31 oppon increasing the ionization potential from 30 to 70 eV, typically the m/e = 2 ion current for the KRI sample increased by a factor of about 1000 under the same pressure conditions as those of the ultrapure hydrogen control. Fig. 31 is representative of the results of the determination of the increase in the m/e = 2 ion current for each sample of MHX or MHMX upon increasing the ionization potential from 30 to 70 eV under the same pressure conditions as those of the ultrapure hydrogen control.

The m/e = 2 ion current for the KHCl, KH KHCO<sub>3</sub>, CsHI, SrHF, and SrHBr sample increased by a factor of about 400, 200, 400, 20, and 50, respectively, under the same pressure conditions as those of the ultrapure hydrogen control.

## 4. Conclusions

The ToF-SIMS, XPS, NMR, and thermal decomposition with analysis by GC, and MS results confirm the identification of novel hydride compounds MHX and MHMX wherein M is the metal, X, is a singly negatively charged anion, and H comprises a novel high binding energy hydride ion. The negative ToF-SIMS spectra of KHI, KH KHCO<sub>3</sub>. RbHF, and SrHF were dominated by the hydride ion peak which identified the structures. The thermal decomposition with mass spectroscopic analysis indicated the minimum hydrogen or dihydrino content. At least H<sup>-</sup>(1/2) was observed to be present in KHX and KH KHCO<sub>3</sub>.

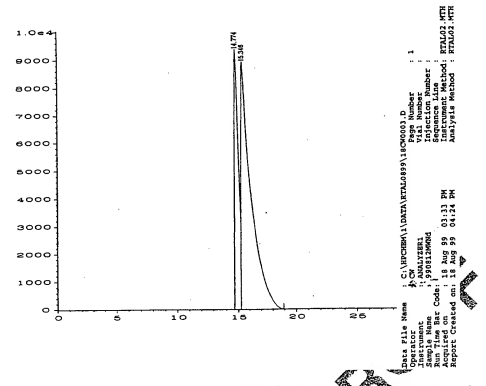


Fig. 29. The gas chromatograph of the dihydrino or hydrogen released from the SrHBr sample when it was heated to above 600°C.

Table 4
The minimum dihydrino or hydrogen content of novel hydride compounds determined by gas chromatograph of the gas released from each sample when it was heated to above 600°C

Novel hydride	Hydrogen or
compound	dihydrino content
•	(μmol/g)
KHI	400
KHCI	200
KHKHCO <sub>3</sub>	4.0
SrHF	169
SrHBr	130
	ATTA TOTAL

The XPS of the low binding energy region of KHI is consistent with the presence of H (n = 1/6)  $E_b = 22.8$  eV. This product is predicted by an autocatalysis reaction of two H(1/4) atoms formed via a potassium catalyst which has been confirmed by extreme ultraviolet spectroscopy [3]. The XPS of KHGI is consistent with the presence of a new paragraph of Eq. (2) eV which is in agreement with the predicted binding energy of  $H^-(n = 1/8)$   $E_b = 36.1$  eV. The bonding of hydrino hydride ions with alkali and alkaline eith metals significantly broadens the metal core level peaks. The magnitude of broadening ranges from 0.3 to 0.9 eV, depending on compound. The

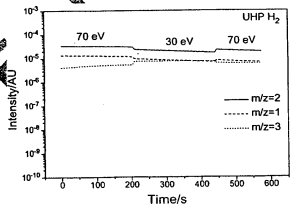


Fig. 30. The intensity as a function of time for masses m/e = 1, m/e = 2, and m/e = 3 obtained while changing the ionization potential (IP) of the mass spectrometer from 30 to 70 eV for ultrapure hydrogen.

trend for the broadening effect follows the sequence: iodo hydrides~bromo hydrides > fluoro hydrides~chloro hydrides. In particular, two additional spin-orbit splittings had to be added to each of potassium iodo hydride, calcium bromo hydride, and strontium bromo hydride in order to obtain a good curve fit of the K 2p, Ca 2p, and Sr 3d spectra, respectively.

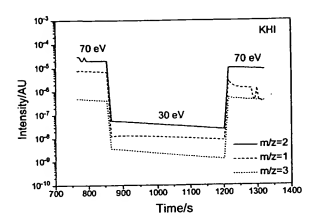


Fig. 31. The intensity as a function of time for masses m/e = 1, m/e = 2, and m/e = 3 obtained while changing the ionization potential (IP) of the mass spectrometer from 30 to 70 eV for gas released from thermal decomposition of the KHI sample.

Large distinct upfield resonances were observed at -3.2 ppm, -4.6 ppm, -4.1 ppm, -4.4 ppm in the case of KHI, KHCI, KHBr, and RbHF, respectively. The peaks are assigned to novel hydride ions that have substantially smaller radii than that of ordinary hydride ion since the shift was extraordinarily far upfield. The NMR peak of KHI at -3.2 ppm may be due to  $H^-(n = 1/6)$   $E_b = 22.8$  eV observed by XPS. The NMR peak of KHCI at -4.6 ppm may be due to  $H^-(n = 1/8)$   $E_b = 36.1$  eV observed by XPS. The NMR of SrHBr showed a  $^1H^{-1}H$  separation of under 1.5 Å . This is very significant compared to the atomic spacing parameters of ordinary strontium hydride and is consistent with a novel hydride ion of a greatly reduced radius.

The chemical structure and properties of the present novel compounds are indicative of a new field of hydrogar chemistry. Novel hydride ions may combine with other cations such as other alkali cations and alkaline early, rare and transition element cations. Numerous not be ompounds may be synthesized with extraordinary properties relative to the corresponding compounds having reliative to the corresponding compounds having reliative to the corresponding compounds may have breath of applications. For example, a high voltage battery according to the hydride binding energies observed by XPS may be possible having projected specifications that surpass those of the internal combustion engine. The discovery of a novel hydride ion with a high binding energy has implications for a new field of hydride themistry with applications such as a high voltage batter 19

voltage batter 19 Hydride tons having extraordinary binding energies may stabilize a ation Mx+ in an extraordinarily high oxidation state such as 42 in the case of lithium. Thus, these hydride ions may be used as the basis of a high voltage battery of a rocking chair design wherein the hydride ion moves back and forth between the cathode and anode half cells

during discharge and charge cycles. Exemplary reactions for a cation M<sup>r+</sup> are:

Cathode reaction:

$$MH_x + e^- \rightarrow MH_{x-1} + H^- \tag{1}$$

Anode reaction:

$$MH_{x-2} + H^- \rightarrow MH_{x-1} + e^-$$
 (2)

Overall reaction:

$$MH_x + MH_{x-2} \to 2MH_{x-1}$$
. (3)

#### Appendix A.

It has been reported that intense extreme ultraviolet (EUV) emission was observed at low temperatures (e.g. ≈ 10³ K) from atomic hydrogen and certain atomized elements or certain gaseous ions which ionize at integer multiples of the potential energy of atomic hydrogen, 27.2 eV [1-6]. The mechanism of EUV emission can not be explained by the conventional chemistry of hydrogen, but it is predicted by a solution of the schrodinger equation with a nonradiative boundary constraint put forward by Mills [12,13]. The energy release must result in a lower-energy state of hydrogen. Mills predicted that certain atoms or ions serve as catalysts to release energy from hydrogen to produce an increased binding energy hydrogen atom called a hydrino atom having a binding energy of

Binding Energy 
$$\frac{13.6 \text{ eV}}{n^2}$$
 (A.1)

$$\frac{1}{4}, \dots, \frac{1}{n} \tag{A.2}$$

where  $a_{\rm H}$  is the radius of the hydrogen atom. Hydrinos are predicted to form by reacting an ordinary hydrogen atom with a catalyst having a net enthalpy of reaction of about

$$m \cdot 27.2 \text{ eV}$$
 (A.3)

where m is an integer. This catalysis releases energy from the hydrogen atom with a commensurate decrease in size of the hydrogen atom,  $r_n = na_H$ . For example, the catalysis of H(n = 1) to H(n = 1/2) releases 40.8 eV, and the hydrogen radius decreases from  $a_H$  to  $\frac{1}{2}a_H$ .

The excited energy states of atomic hydrogen are also given by Eq. (A.1) except that

$$n = 1, 2, 3, \dots$$
 (A.4)

The n=1 state is the "ground" state for "pure" photon transitions (the n=1 state can absorb a photon and go to an excited electronic state, but it cannot release a photon and go to a lower-energy electronic state). However, an electron transition from the ground state to a lower-energy state is possible by a nonradiative energy transfer such as multipole coupling or a resonant collision mechanism. These

lower-energy states have fractional quantum numbers, n =1/integer. Processes that occur without photons and that require collisions are common. For example, the exothermic chemical reaction of H + H to form H<sub>2</sub> does not occur with the emission of a photon. Rather, the reaction requires a collision with a third body, M, to remove the bond energy-H + H +  $M \rightarrow$  H<sub>2</sub> + M\* [14]. The third body distributes the energy from the exothermic reaction, and the end result is the H2 molecule and an increase in the temperature of the system. Some commercial phosphors are based on nonradiative energy transfer involving multipole coupling. For example, the strong absorption strength of Sb3+ ions along with the efficient nonradiative transfer of excitation from Sb3+ to Mn2+, are responsible for the strong manganese luminescence from phosphors containing these ions [15]. Similarly, the n = 1 state of hydrogen and the n=1/integer states of hydrogen are nonradiative, but a transition between two nonradiative states is possible via a nonradiative energy transfer, say n=1 to n=1/2. In these cases, during the transition the electron couples to another electron transition, electron transfer reaction, or inelastic scattering reaction which can absorb the exact amount of energy that must be removed from the hydrogen atom. Thus, a catalyst provides a net positive enthalpy of reaction of  $m \cdot 27.2$  eV (i.e. it absorbs  $m \cdot 27.2$  eV where m is an integer). Certain atoms or ions serve as catalysts which resonantly accept energy from hydrogen atoms and release the energy to the surroundings to effect electronic transitions to fractional quantum energy levels.

#### A.1. Inorganic catalysts

A catalytic system is provided by the ionization of t electrons from an atom to a continuum energy level such that the sum of the ionization energies of the t electrons is approximately  $m \cdot 27.2$  eV where m is an integer. One such catalytic system involves strontium. The first through the fifth ionization energies of strontium are 5.69484, 11.0301, 42.8957, and 71.6 eV, respectively [16]. The ionization reaction of Sr to Sr<sup>5+</sup> (t = 5), then, has a net enthalphy feaction of 188.2 eV, which is equivalent to m = 24.024.

188.2 eV + Sr(m) + H 
$$\left[\frac{a_{\rm H}}{p}\right]$$
  $\rightarrow$  5.4  
+5e<sup>-</sup> + H  $\left[\frac{a_{\rm H}}{(p+7)}\right]$  + [(p 7)<sup>2</sup> - p<sup>2</sup>]Y 13.6 eV, (A.5)

$$Sr^{5+} + 5e^{-} \rightarrow Sr(m) + 188.2 \text{ eV}$$
 (A.6)

And, the overall reaction is

H 
$$\left[\frac{a_{\rm H}}{p}\right]$$
 +  $\left[(p+7)^2 - p^2\right]X13.6$  eV. (A.7)

Another catalytic system that is provided by the ionization of t electrons from an atom to a continuum energy level such

that the sum of the ionization energies of the t electrons is approximately mX27.2 eV where m is an integer involves potassium. The first, second, and third ionization energies of potassium are 4.34066, 31.63, 45.806 eV, respectively [16]. The triple ionization (t=3) reaction of K to,  $K^{3+}$  then, has a net enthalpy of reaction of 81.7426 eV, which is equivalent to m=3 in Eq. (A.3).

81.7426 eV + K(m) + H 
$$\left[\frac{a_{\rm H}}{p}\right] \rightarrow K^{3+} + 3e^{-}$$
  
+H  $\left[\frac{a_{\rm H}}{(p+3)}\right]$  +  $[(p+3)^2 - p^2]X13.6$  eV, (A.8)

$$K^{3+} + 3e^{-} \rightarrow K(m) + 81.7426 \text{ eV}.$$
 (A.9)

And, the overall reaction is

$$H\left[\frac{a_{H}}{p}\right] \to H\left[\frac{a_{H}}{(p+3)}\right] + [(p+3)^{2} - p^{2}]X13.6 \text{ eV}.$$
(A.10)

A novel hydride ion having extraordinal schemical properties given by Mills [12] is predicted to form by the reaction of an electron with a hydrine (Eq. (A.11)). The resulting hydride ion is referred to as a hydrino hydride ion, designated as  $H^{-}(1/p)$ .

$$H\left[\frac{a_{\rm H}}{p}\right] + e^- \to 0$$
 (A.11)

The hydrific hydride ion is distinguished from an ordinary hydride ion having a binding energy of 0.8 eV. The hydring hydride ion is predicted [12] to comprise a hydrogen nucleus and two indistinguishable electrons at a binding energy according to the following formula:

Binding Energy = 
$$\frac{\hbar^2 \sqrt{s(s+1)}}{8\mu_e a_0^2 \left[\frac{1+\sqrt{s(s+1)}}{p}\right]^2} - \frac{\pi\mu_0 e^2 \hbar^2}{m_e^2 a_0^3} \left(1 + \frac{2^2}{\left[\frac{1+\sqrt{s(s+1)}}{p}\right]^3}\right)$$
(A.12)

where p is an integer greater than one, s = 1/2,  $\pi$  is pi,  $\hbar$  is Planck's constant bar,  $\mu_0$  is the permeability of vacuum,  $m_e$  is the mass of the electron,  $\mu_e$  is the reduced electron mass,  $a_0$  is the Bohr radius, and e is the elementary charge. The ionic radius is

$$r_1 = \frac{a_0}{n}(1 + \sqrt{s(s+1)}); \quad s = \frac{1}{2}$$
 (A.13)

From Eq. (A.13), the radius of the hydrino hydride ion  $H^-(1/p)$ ; p = integer is 1/p that of ordinary hydride ion,  $H^-(1/1)$ . Compounds containing hydrino hydride ions have been isolated as products of the reaction of atomic hydrogen with atoms and ions identified as catalysts by EUV emission [1-6,17-20].

Alkali and alkaline earth hydrides react violently with water to release hydrogen gas which subsequently ignites due to the exothermic reaction with water. Typically metal hydrides decompose upon heating at a temperature well below the melting point of the parent metal. These saline hydrides, so called because of their saltlike or ionic character, are the monohydrides of the alkali metals and the dihydrides of the alkaline-earth metals. Mills predicts a hydrogen-type molecule having a first binding energy of about

Binding Energy = 
$$\frac{15.5}{(\frac{1}{p})^2}$$
 eV. (A.14)

Dihydrino molecules may be produced by the thermal decomposition of hydrino hydride ions. KH-(1/2) may be less reactive and more thermally stable than ordinary potassium hydride, but may react at high temperature to form a hydrogen-type molecule. For example, potassium iodo hydride KH(1/2)I may be heated to release dihydrino by thermal decomposition.

$$2KH(1/2)I \xrightarrow{\Delta} H_2^* \left[2c' = \frac{a_0}{\sqrt{2}}\right] + 2KI$$
 (A.15)

where 2c' is the internuclear distance and  $a_0$  is the Bohr radius [12]. The possibility of releasing dihydrino by thermally decomposing alkali and alkaline earth halido hydrides and KH KHCO3 with identification by gas chromatography was explored.

The first ionization energy, IP1, of the dihydrino molecule

$$H_2^* \left[ 2c' = \frac{\sqrt{2}a_0}{2} \right] \to H_2^* [2c' = a_0]^+ + e^-$$
 (A.16)

is  $1P_1=62$  eV(p=2 in Eq. (A.14)); whereas, the first ionization energy of ordinary molecular hydrogen,  $H_2[2c'=\sqrt{2}a_0]$ , is 15.46 eV. Thus, the possibility of using mass spectroscopy to discriminate  $H_2[2c'=\sqrt{2}a_0]$  from  $H_2^*[2c'=a_0/\sqrt{2}]$  on the basis of the large difference between the ionization energies of the two species was explored. A novel high binding energy hydrogen molecule assigned to dihydrino H2 [2c/  $a_0/\sqrt{2}$ ] was identified by the thermal decomposition of alkali and alkaline earth halido hydrides and KH KH@ analysis by gas chromatography, and mass spectros

#### References

[1] Mills R, Dong J, Lu, Y. Observation of extreme ultraviolet hydrogen emission from incandescently heated hydrogen gas with certain catalysts. Pacific Conference on Chemistry and Spectroscopy and the Silve CS Vestern Regional Meeting, Ontario Convention enter, California, October 6-8, 1999.

- [2] Mills R, Dong J, Lu Y. Observation of extreme ultraviolet hydrogen emission from incandescently heated hydrogen gas with certain catalysts. Int J Hydrogen Energy 2000;25:919-43.
- [3] Mills R. Observation of extreme ultraviolet emission from hydrogen-KI plasmas produced by a hollow cathode discharge. Int J Hydrogen Energy, submitted for publication.
- [4] Mills R. Temporal behavior of light-emission in the visible spectral range from a Ti-K2CO3-H-Cell. Int J Hydrogen Energy, accepted.
- [5] Mills R, Lu R, Onuma T. Formation of a hydrogen plasma from an incandescently heated hydrogen-potassium gas mixture and plasma decay upon removal of heater power. Int J Hydrogen Energy, accepted.
- [6] Mills R, Nansteel M, Lu Y. Observation of extreme ultraviolet hydrogen emission from incandescently heated hydrogen gas with strontium that produced an optically measured power balance that was 4000 times the control. Int J Hydrogen Energy, accepted.
- [7] Microsc. Microanal. Microstruct., 3(1) 1992
- [8] PHI Trift II, ToF-SIMS Technical Broggare, Eden Prairie, MN 55344, USA, 1999.
- MN 55544, Oct., 17. Hydrogen Energy 2000;25:669-83.
- [10] Briggs D, Seah MP, editors. Practical surface analysis, 2nd ed.
  lon and neutral spectroscopy to ... New York: Wiley, 1992.
  [11] Mackay K.M. Hydroger Compounds of the metallic elements.
  London: E. & F. N. Spon, 1966.
  [12] Mills R. The grand unified theory of classical quantum
- Mechanics. Cranburg Ny BlackLight Power, Inc. January 2000. Distributed by Amazon.com.
- [13] Mills R the hidrogen atom revisited. Int J Hydrogen Energy, accepted.
- [14] Sidewick NV The chemical elements and their compounds, ol. 12 Oxford: Clarendon Press, 1950. p. 17.
- tamb MD. Luminescence spectroscopy. London: Academic P 978. p. 68.
- Linde DR. CRC handbook of chemistry and physics, 79th ed. Boca Raton, FL: CRC Press, (1998-9). pp. 10-175-10-177.
- [N] Mills R, Dhandapani B, Greenig N, He J. Synthesis and characterization of potassium iodo hydride. Int J Hydrogen Energy, accepted.
- [18] Mills R. Novel hydrogen compounds from a potassium carbonate electrolytic cell. Fusion Technol 2000;37(2):157-
- [19] Mills R, He J, Dhandapani B. Novel hydrogen compounds. 1999 Pacific Conference on Chemistry and Spectroscopy and the 35th ACS Western Regional Meeting, Ontario Convention Center, California, October 6-8, 1999.
- [20] Mills R. Highly stable novel inorganic hydrides. J Mater Res, submitted for publication.

THIS PAGE BLANK (USPTO)